STN-Structure seaseh 5.23.06 10/613,650

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ANSWER 1 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:1154157 CAPLUS

DOCUMENT NUMBER:

143:422465

TITLE:

Preparation of phosphonate analogs of HIV protease inhibitors and methods for identifying anti-HIV

therapeutic compounds

INVENTOR(S):

Arimilli, Murty N.; Becker, Mark M.; Birkus, Gabriel

PATENT ASSIGNEE(S):

SOURCE:

U.S. Pat. Appl. Publ., 1034 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

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PRIORITY APPLN. INFO.:
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WO 2003-US12943

US 2003-740694

A2 20030425

A 20031222

GI

The invention relates to phosphonate-substituted carbamates I and cyclic ureas II [wherein A = A1, A2, or W3 with the proviso that at least one of A = A1; A1 = [Y2(CR2R2)1-12]0-12Y2W6; A2 = [Y2(CR2R2)1-12]0-12Y2W3; W3 = substituted (hetero)cyclyl, R5, C(Y1)R5, C(Y1)W5, SO2R5, or SO2W5; W5 = substituted (hetero)cyclyl; W6 = triphosphono-substituted W3; Y1 = O, S, N(Rx), N(O)(Rx), N(ORx), N(O)(ORx), or N(N(Rx)2); Y2 = independently a bond, O, N(Rx), N(O)(Rx), N(O)(ORx), N(O)(ORx), N(N(Rx)2), SO0-2, or SO0-2SO0-2; Rx = independently H, R1, W3, a protecting group, etc.; R1 = independently H or alkyl; R2 = independently H, R1, halo, CN, N3, NO2, Y1, Rx, N(Rx)2, SO-2Rx, substituted alkyl, alkenyl, alkynyl, etc.; R3 = halo, CN, N3, NO2, Y1, Rx, N(Rx)2, SRx, SORx, SO2Rx, OC(Y1)Rx, OC(Y1)ORx, C(Y1)Rx, etc. with provisos; R5 = substituted alkyl, alkenyl, or alkynyl;

RN 622865-50-3 CAPLUS

CN 10,13-Dioxa-2,6-diaza-12-phosphapentadecanoic acid, 12-ethoxy-4-hydroxy-6-[(4-methoxyphenyl)sulfonyl]-8,8-dimethyl-3-(phenylmethyl)-, 1,1-dimethylethyl ester, 12-oxide, (3S,4R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 2 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

2005:612479 CAPLUS 143:97530

TITLE:

Preparation of phosphonate analogs of HIV protease inhibitors and methods for identifying anti-HIV

therapeutic compounds

INVENTOR (S):

Arimilli, Murty N.; Becker, Mark M.; Birkus, Gabriel; Bryant, Clifford; Chen, James M.; Chen, Xiaowu; Cihlar, Tomas; Dastgah, Azar; Eisenberg, Eugene J.; Fardis, Maria; Hatada, Marcos; He, Gong-Xin; Jin, Haolun; Kim, Choung U.; Lee, William A.; Lee, Christopher B.; Lin, Kuci Ving, Lin, Hongton, Mackey, Markey, Markey

Haolun; Kim, Choung U.; Lee, William A.; Lee, Christopher P.; Lin, Kuei-Ying; Liu, Hongtao; Mackman, Richard L.; McDermott, Martin J.; Mitchell, Michael L.; Nelson, Peter H.; Pyun, Hyung-Jung; Rowe, Tanisha D.; Sparacino, Mark; Swaminathan, Sundaramoorthi; Tario, James D.; Wang, Jianying; Williams, Matthew A.; Xu, Lianhong; Yang, Zheng-Yu; Yu, Richard H.; Zhang,

Jiancun; Zhang, Lijun

PATENT ASSIGNEE(S):

SOURCE:

Gilead Sciences, Inc., USA PCT Int. Appl., 1723 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

8

PATENT INFORMATION:

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US 2005	MR, NE	-	•		2005:	1027	,	TC 2	002	7400	0.4		_	0001	200
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GΙ

The invention relates to phosphonate-substituted carbamates I and cyclic ureas II [wherein A = A1, A2, or W3 with the proviso that at least one of A = A1; A1 = [Y2(CR2R2)1-12]0-12Y2W6; A2 = [Y2(CR2R2)1-12]0-12Y2W3; W3 = substituted (hetero)cyclyl, R5, C(Y1)R5, C(Y1)W5, SO2R5, or SO2W5; W5 =

10/613,650

RN 622865-50-3 CAPLUS

CN 10,13-Dioxa-2,6-diaza-12-phosphapentadecanoic acid, 12-ethoxy-4-hydroxy-6-[(4-methoxyphenyl)sulfonyl]-8,8-dimethyl-3-(phenylmethyl)-, 1,1-dimethylethyl ester, 12-oxide, (3S,4R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 3 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:99287 CAPLUS

DOCUMENT NUMBER: 140:339141

TITLE: Novel arylsulfonamides possessing sub-picomolar HIV

protease activities and potent anti-HIV activity against wild-type and drug-resistant viral strains Miller, John F.; Furfine, Eric S.; Hanlon, Mary H.;

Hazen, Richard J.; Ray, John A.; Robinson, Laurence; Samano, Vicente; Spaltenstein, Andrew

CORPORATE SOURCE: GlaxoSmithKline, Research Triangle Park, NC, 27709,

USA

SOURCE: Bioorganic & Medicinal Chemistry Letters (2004),

14(4), 959-963

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 140:339141

GI

AUTHOR (S):

Furanofuryl analogs of the HIV protease inhibitor amprenavir such as I are prepared in which a terminally substituted n-alkyl group is appended to the N-iso-Bu group of amprenavir and in which the substituents on the N-arylsulfonyl moiety are varied. Some of the inhibitors such as I are found to have greatly enhanced inhibition of HIV protease; the amprenavir analogs also inhibit the growth of both wild-type and resistant strains of HIV and are more effective against the HIV strains than the currently marketed HIV protease inhibitors amprenavir, indinavir, and nelfinavir. E.g., I inhibits wild-type HIV protease with a Ki value of 0.014 pM, and inhibits wild-type and resistant strains of HIV with IC50 values of between 1.6 nM and 15 nM.

IT 681028-81-9P

CN

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of furanofuryl amprenavir analogs with modifications at the N-arylsulfonyl and N-iso-Bu moieties which show improved HIV protease inhibition and inhibition of wild-type and resistant HIV strains)

Ι

RN 681028-81-9 CAPLUS

Carbamic acid, [(1S,2R)-3-[(1,3-benzodioxol-5-ylsulfonyl)(5-cyano-2,2-dimethylpentyl)amino]-2-hydroxy-1-(phenylmethyl)propyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 4 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:875072 CAPLUS

DOCUMENT NUMBER: 139:381610

TITLE: Preparation of phosphonate analogs of HIV protease

inhibitors and methods for identifying anti-HIV

therapeutic compounds

INVENTOR(S): Birkus, Gabriel; Chen, James M.; Chen, Xiaowu; Cihlar,

Tomas; Eisenberg, Eugene J.; Hatada, Marcos; He, Gong-Xin; Kim, Choung U.; Lee, William A.; McDermott,

APPLICATION NO.

DATE

Martin J.; Swaminathan, Sundaramoorthi

PATENT ASSIGNEE(S): Gilead Sciences, Inc., USA SOURCE: PCT Int. Appl., 814 pp.

KIND

CODEN: PIXXD2

DATE

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.

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L4 ANSWER 5 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2003:875071 CAPLUS 139:381609

DOCUMENT NUMBER: TITLE:

Preparation of phosphonate analogs of HIV protease inhibitors with improved cellular accumulation

properties

INVENTOR (S):

Arimilli, Murty N.; Becker, Mark M.; Bryant, Clifford; Chen, James M.; Chen, Xiaowu; Dastgah, Azar; Fardis, Maria; He, Gong-Xin; Jin, Haolun; Kim, Choung U.; Lee, William A.; Lee, Christopher P.; Lin, Kuei-Ying; Liu, Hongtao; Mackman, Richard L.; Mitchell, Michael L.; Nelson, Peter H.; Pyun, Hyung-Jung; Rowe, Tanisha D.; Sparacino, Mark; Swaminathan, Sundaramoorthi; Tario, James D.; Wang, Jianying; Williams, Matthew A.; Xu, Lianhong; Yang, Zheng-Yu; Yu, Richard H.; Zhang,

PATENT ASSIGNEE(S):

SOURCE:

Jiancun; Zhang, Lijun Gilead Sciences, Inc., USA PCT Int. Appl., 1727 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
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AU 2003231765	A1 20031110	AU 2003-231765	20030425
BR 2003009573	A 20050201	BR 2003-9573	20030425
EP 1509537	A2 20050302	EP 2003-747326	20030425
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               PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
          RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
              BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     AU 2003291998
                              A1
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                                                                                20031106
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                                                                               20031106
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, FI, RO, CY, TR, BG, CZ, EE, HU, SK
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PRIORITY APPLN. INFO.:
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                                                    US 2003-423496
                                                                            A2 20030425
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                                                    US 2003-465810P
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                                                                            Α
                                                                               20030425
                                                    WO 2003-EP12423
                                                                            W 20031106
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OTHER SOURCE(S): MARPAT 139:381609

AB The invention relates to phosphonate-substituted carbamates I and cyclic ureas II [wherein A = A1, A2, or W3 with the proviso that at least one of

## Absolute stereochemistry.

RN 622865-50-3 CAPLUS

CN 10,13-Dioxa-2,6-diaza-12-phosphapentadecanoic acid, 12-ethoxy-4-hydroxy-6-[(4-methoxyphenyl)sulfonyl]-8,8-dimethyl-3-(phenylmethyl)-, 1,1-dimethylethyl ester, 12-oxide, (3S,4R)- (9CI) (CA INDEX NAME)

## Absolute stereochemistry.

ANSWER 6 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2002:814117 CAPLUS

DOCUMENT NUMBER:

137:325410

TITLE:

Broad-spectrum 2-(substituted-amino)-

benzothiazolesulfonamide HIV protease inhibitors

INVENTOR (S): Surleraux, Dominique Louis Nestor Ghislain; Wigerinck,

Piet Tom Bert Paul; Getman, Daniel; Verschueren, Wim

Gaston; Vendeville, Sandrine; De Bethune,

Marie-Pierre; De Kerpel, Jan Octaaf Antoon; Moors, Samuel Leo Christiaan; De Kock, Herman Augustinus;

Voets, Marieke Christiane Johanna

PATENT ASSIGNEE(S):

Tibotec Pharmaceuticals Ltd., Ire.

SOURCE:

PCT Int. Appl., 83 pp.

DOCUMENT TYPE:

CODEN: PIXXD2

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE heterocyclyloxyalkyl, aralkyl, aryloxyalkyl, (un)substituted aminoalkyl; R5NAR6 = heterocyclic) their N-oxides, salts, stereoisomeric forms, racemic mixts., prodrugs, esters and metabolites were prepared I are useful as broad-spectrum HIV protease inhibitors, and may be formulated in diagnostic kits. Thus, the sulfonamide II, prepared in several steps from the benzothiazolecarbamate, showed activity against a number of resistant mutants of HIV-1 strain LAI.

473739-04-7P TT

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(broad-spectrum 2-aminobenzothiazolesulfonamide HIV protease inhibitors)

RN 473739-04-7 CAPLUS

CN Carbamic acid, [(1S,2R)-3-[[[2-(acetylamino)-6-benzothiazolyl]sulfonyl][2-(2-pyridinylamino)ethyl]amino]-2-hydroxy-1-(phenylmethyl)propyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 7 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2000:900607 CAPLUS

DOCUMENT NUMBER:

134:56676

TITLE:

Preparation of arylsulfonamides as inhibitors of

aspartyl protease

INVENTOR (S):

Hale, Michael Robin; Tung, Roger; Price, Stephen; Wilkes, Robin David; Schairer, Wayne Carl; Jarvis, Ashley Nicholas; Spaltenstein, Andrew; Furfine, Eric Steven; Samano, Vicente; Kaldor, Istvan; Miller, John

Franklin; Brieger, Michael Stephen

PATENT ASSIGNEE(S):

Vertex Pharmaceuticals Inc., USA; et al.

SOURCE:

PCT Int. Appl., 396 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

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WO 2000	NO 2000076961 W: AE, AG, AI				-	2000	 1221	,	WO 2	 000-1	US15	781		2	0000	608
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ŧ	ID,	IL,	IN,	IS,	JP,	KE, MN,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,

IT

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SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU,
             ZA, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
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             CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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                                 20020410
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     EP 1194404
                                 20060503
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PRIORITY APPLN. INFO.:
                                             US 1999-139070P
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                                             US 2000-190211P
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                                                                     20000317
                                             WO 2000-US15781
                                                                  W 20000608
                                             US 2000-591464
                                                                  A3 20000609
OTHER SOURCE(S):
                         MARPAT 134:56676
```

The title arylsulfonamides, namely (3R,3aS,6aR)-hexahydrofuro[2,3-b]furan-3-yl 3-arylsulfonylamino-1-(4-hydroxyphenyl)-2-hydroxypropylcarbamate derivs. (e.g. I) are prepared These compds. are particularly well suited for inhibiting HIV-1 and HIV-2 protease activity and consequently, may be advantageously used as anti-viral agents against the HIV-1 and HIV-2 viruses. They are useful for treating with a patient diagnosed with AIDS, AIDS related complex (ARC), progressive generalized lymphadenopathy (PGL), Kaposi's sarcoma, thrombocytopenic purpura, or AIDS-related neurol. conditions such as AIDS dementia complex, multiple sclerosis or tropical paraperesis, etc. Thus, (3R,3aS,6aR)-hexahydrofuro[2,3-b]furan-3-yl 3-[N-(1,3-benzodioxol-5-ylsulfonyl)-N-isobutylamino]-1-(4-hydroxyphenyl)-2-hydroxypropylcarbamate underwent Mitsunobu reaction with phenethyl alc. using Ph3P and di-tert-Bu azodicarbonate in CH2Cl2 at room temperature for 1.5

to give 72% I. I showed IC50 of <0.001, <0.001, and 0.01-0.001  $\mu M$  against drug-resistant HIV strains, i.e. wild type, mutant HIV-1 EP13, and mutant D545701-14 HIV strains, resp., in MT-4 cells.

313683-12-4P 313683-13-5P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological

RN 313681-96-8 CAPLUS

CN 4-Oxa-2,10,14-triazapentadecan-15-oic acid, 10-(1,3-benzodioxol-5-ylsulfonyl)-12-hydroxy-8,8-dimethyl-3-oxo-13-[[4-(phenylmethoxy)phenyl]methyl]-, 1,1-dimethylethyl ester, (12R,13S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENÇE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LA ANSWER 8 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

7

ACCESSION NUMBER: 2000

2000:573770 CAPLUS

DOCUMENT NUMBER:

133:177157

TITLE:

Preparation of [1-benzyl-2-hydroxy-3-

(arylsulfonamido)propyl]carbamates as HIV aspartyl

protease inhibitors

INVENTOR(S):

Hale, Michael R.; Baker, Christopher T.; Stammers, Timothy A.; Sherrill, Ronald G.; Spaltenstein, Andrew; Furfine, Eric S.; Maltais, Francois; Andrews, Clarence

Webster, III; Miller, John F.; Samano, Vicente

PATENT ASSIGNEE(S):

Vertex Pharmaceuticals Incorporated, USA

SOURCE:

PCT Int. Appl., 369 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

GT

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PRIORIT	II APP	LIN.	INFO	. :							L999-						
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									1	US 2	2001-	9272	71		A3 2	0010	809
OTHER S	SOURCE	(S):			MARI	TAS	133:	17715	57								

AB 
ABxN(Gx)CH(D)CH(OR7)CH2ND'E'E [wherein A = H, or (un)substituted Ht, R1Ht, or R1Ak; Ak = alkyl; Ht = cycloalkyl, cycloalkenyl, or (un)substituted aryl or heterocyclyl; R1 = CO(CO), (O)SO2, O2C, or (un)substituted NHSO2 or NHCO(CO); B = (un)substituted NHCH2CO; x = 0 or 1; G = H, R7, alkyl; or G may be bound to R7 to form a heterocyclic ring; R7 = H, (CH2O)xY(ZM)(:X)Z(M)x; etc.; M = H, Li, Na, K, Mg, Ca, Ba, alkyl, alkenyl, etc.; X = O or S; Y = P or S; Z = H, O, S, or (un)substituted NH2; D = independently Q or (un)substituted (cyclo)alkyl or (cyclo)alkenyl; Q = (un)substituted carbocylyl or heterocyclyl; D' = (un)substituted alkyl, alkenyl; alkynyl; E = Ht, OHt, HtHt, alkoxy, (un)substituted NH2, alkyl, or carbocyclyl; E' = CO or SO2] were prepared as antiviral agents against HIV-1 and HIV-2 viruses. Thus, 3-NO2C6H4SO2Cl was added to tert-Bu (1S,2R)-N-[1-benzyl-3-[(4-cyano-2,2-dimethylbutyl)amino]-2-hydroxypropyl]carbamate (preparation given) to form the 3-

Ι

ANSWER 9 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2000:304314 CAPLUS

DOCUMENT NUMBER:

132:322147

TITLE:

Preparation of  $\alpha$ - and  $\beta$ -amino acid

hydroxyethylamino sulfonamides as retro viral protease

inhibitors.

INVENTOR(S):

Vazquez, Michael L.; Mueller, Richard A.; Talley, John J.; Getman, Daniel P.; Decrescenzo, Gary A.; Freskos, John N.; Heintz, Robert M.; Bertenshaw, Deborah E.

PATENT ASSIGNEE(S):

SOURCE:

G.D.Searle and Co., USA U.S., 93 pp., Cont.-in-part of Appl. PCT/US93/07814. CODEN: USXXAM

DOCUMENT TYPE:

Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

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	62487				В1	2	2001	0619	1	US 1	L999-	2880	80		19	99904	108	
	65008	32			B1	2	2002	1231	1	US 2	2000-	5251	61		20	0000	314	
US	20020	5239	99		A1	2	2002	0502	1	US 2	2001-	7982	55		20	0103	305	

US 6417387	B2	20020709				
US 2003191319	<b>A1</b>	20031009	US	2002-157019		20020530
US 6646010	B2	20031111				
US 2004044047	A1	20040304	US	2002-199481		20020722
US 6846954	B2	20050125				
US 6924286	B1	20050802	US	2003-633376		20030804
US 2004229922	A1	20041118	US	2004-812343		20040330
US 2005267171	A1	20051201	US	2005-110943		20050421
PRIORITY APPLN. INFO.:			US	1992-934984	B2	19920825
			WO	1993-US7814	A2	19930824
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			US	1995-451090	A3	19950525
			US	1999-288080	A1	19990408
			US	2001-798255	A1	20010305
			US	2002-157019	A1	20020530
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			US	2003-633376	A1	20030804
OMITTED COLIDGE (C)	1/3 D D 3 C					

OTHER SOURCE(S): MARPAT 132:322147

GI

Amino acid hydroxyethylamino sulfonamide compds. I [R2 = (un)substituted aryl, (cyclo)alkyl, aralkyl, cycloalkylalkyl; R3 = alkyl, haloalkyl, alkenyl, alkynyl, hydroxy-, alkoxy-, alkylthio-, or alkylsulfonylalkyl, cycloalkylalkyl, heterocycloalkyl, heterocycloalkylalkyl, aryl, aralkyl, or heteroaralkyl; R4 = heterocycloalkyl, heteroaryl or aryl; Y = O or S; A = heterocycloalkyl, heterocycloalkoxy, heterocycloalkylalkoxy, heterocycloalkylalkoxy, heteroaralkyl, heteroarylalkoxy, heteroaryloxy or heteroaryl] were prepared as retroviral protease inhibitors, particular as inhibitors of HIV protease. Thus, compound II (Cbz = benzyloxycarbonyl) was prepared and assayed for HIV inhibitory activity (IC50 = 16 nM). Compds. of formula I were tested for cytotoxicity and efficacy (IC50, EC50 and TD50 values at the nanomolar level are tabulated).

IT 169281-15-6P 169281-16-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(amino acid hydroxyethylamino sulfonamides as retroviral protease inhibitors)

RN 169281-15-6 CAPLUS

CN Carbamic acid, [(1S,2R)-2-hydroxy-3-[(3-methoxypropyl) (phenylsulfonyl)amin o]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 169281-16-7 CAPLUS

CN Carbamic acid, [(1S,2R)-2-hydroxy-3-[[3-(methylthio)propyl](phenylsulfonyl)amino]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 10 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:220728 CAPLUS

DOCUMENT NUMBER: 132:265504

TITLE: Preparation of hydroxyethylamino sulfonamides useful

as retroviral protease inhibitors.

INVENTOR(S): Vazquez, Michael L.; Mueller, Richard A.; Talley, John

J.; Getman, Daniel P.; Decrescenzo, Gary A.; Freskos,

John N.; Bertebshaw, Deborah E.; Heintz, Robert M.

PATENT ASSIGNEE(S): G.D. Searle and Co., USA

SOURCE: U.S., 119 pp., Cont.-in-part of U.S. 204,872,

abandoned.
CODEN: USXXAM

CODEN: USXXA

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 6

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6046190 WO 9404492	A A1	20000404 19940303	US 1996-586866 WO 1993-US7814	19960124 19930824
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KP, KR, KZ, LK, LU, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD,
               SE, SK, UA, US, VN
          RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
               BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
     EP 810209
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                             Α3
                                    19981202
     EP 810209
                             B1
                                    20020605
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                                                                           19940823
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PRIORITY APPLN. INFO.:
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OTHER SOURCE(S):
                            MARPAT 132:265504
     Hydroxyethylamino sulfonamide compds. R9R10N(CR7R8)pCHR1C(:Y)NR6CHR2CH(OH)
     CH2NR3S(:0) \times R4 [I: R1 = H, CH2SO2NH2, CH2CO2CH3, alkyl, haloalkyl,
     alkenyl, alkynyl, cycloalkyl, amino acid side chains, etc.; R2 =
     (un)substituted alkyl, aryl, cycloalkyl, cycloalkylalkyl, aralkyl; R3 = H,
alkyl, haloalkyl, alkenyl, alkynyl, aryl, heteroaryl, mono- and
     disubstituted aminoalkyl, etc.; R4 = alkyl, haloalkyl, alkenyl, alkynyl,
     aryl, (un)saturated heterocycle, (un)substituted aromatic heterocycloalkyl,
etc.;
     R6 = H, alkyl; Y = O, S, NR3; R7,R8 = independently H, R1, or together
     with R1 and the carbon atoms to which they are attached represent a
     cycloalkyl radical; R9 = H, R3, or R3SO2; R10 = H, alkoxycarbonyl,
     alkylcarbonyl, aroyl, aryloxycarbonyl, heterocyclylalkoxycarbonyl, mono-
     and disubstituted aminocarbonyl, or aminoalkanoyl, etc.; or R9R10N =
     heterocycloalkyl or heteroaryl; x = 0-2; p = 0-1] or their
     pharmaceutically acceptable salts, prodrugs, or esters were prepared as
     inhibitors of retroviral proteases such as human immunodeficiency virus
      (HIV). Many inhibitors were prepared by (1) preparing an N-protected amino
     epoxide and (2) reacting this with an amine and (3) preparing a sulfonamide
     by reacting with a sulfonyl chloride or sulfonyl anhydride in the presence
     of an acid scavenger. The amino function of the sulfonamide was then (4)
     deprotected and (5) reacted with a carboxylate.
     N1-[2R-hydroxy-3-[(3-methylbutyl)(phenylsulfonyl)amino]-1S-
      (phenylmethyl)propyl]-2S-[(2-quinolinylcarbonyl)amino]butanediamide was
     prepared and assayed for HIV protease inhibitory activity (IC50 = 1.5 nM).
     Compds. of formula I were tested for cytotoxicity and antiviral efficacy
      (IC50, EC50, and TD50 values at the nanomolar level are tabulated).
TT
     169281-15-6P 169281-16-7P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
         (preparation of hydroxyethylamino sulfonamides useful as retroviral protease
         inhibitors)
RN
     169281-15-6 CAPLUS
CN
     Carbamic acid, [(1S,2R)-2-hydroxy-3-[(3-methoxypropyl)(phenylsulfonyl)amin
     o]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)
```

Absolute stereochemistry.

RN 169281-16-7 CAPLUS

CN Carbamic acid, [(1S,2R)-2-hydroxy-3-[[3-(methylthio)propyl](phenylsulfonyl
) amino]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX
NAME)

Absolute stereochemistry.

REFERENCE COUNT: 45 THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 11 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:670116 CAPLUS

DOCUMENT NUMBER: 131:295568

TITLE:  $\alpha$ - and  $\beta$ -Amino acid hydroxyethylamino

sulfonamides useful as retroviral protease inhibitors
INVENTOR(S):

Vazques, Michael L.; Mueller, Richard A.; Talley, John
J.; Getman, Daniel P.; Decrescenzo, Gary A.; Freskos,

John N.; Bertenshaw, Deborah E.; Heintz, Robert M.

PATENT ASSIGNEE(S): G. D. Searle and Co., USA

SOURCE: U.S., 130 pp., Cont.-in-part of U.S. 204,827.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 6

PAT	TENT	NO.			KIN	D -	DATE			APPL	ICAT	ION :	NO.		D	ATE	
	5968						1999				994-:					9940	
WO	9404				A1		1994	0303		WO 1	993-1	US78:	14		1:	9930	324
	W:	ΑT,	AU,	BB,	BG,	BR,	BY,	CA,	CH,	CZ,	DE,	DK,	ES,	FI,	GB,	HU,	JP,
		KΡ,	KR,	KZ,	LK,	LU,	MG,	MN,	MW,	NL,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,
		SE,	SK,	UA,	US,	VN											
	RW:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG		•
EP	8102	09			A2		1997	1203		EP 1	997-:	1134	34	•	19	99308	324

		_																	
EP	81020	9			<b>A3</b>		1998:	1202											
EP	81020	9			B1		2002	0605											
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GF	l, IT	', L	I,	LU,	NL,	SE	Ξ,	PT,	ΙE
US	60604	76			Α		2000	0509	τ	JS	1994	-20	482	27			19	940	302
US	62487	75			B1	:	2001	0619	Ţ	JS	1999	-28	808	30			19	990	408
US	20020	5239	9		A1	:	2002	0502	Ţ	JS	2001	-79	825	55			20	010	305
US	64173	87			B2	:	2002	0709											
US	20031	9131	L 9		<b>A1</b>	2	2003	1009	τ	JS	2002	-15	701	<b>.</b> 9			20	020	530
ບຣ	66460	10			B2	2	2003	1111											
ບຣ	69242	86			B1	2	2005	0802	ζ	JS	2003	-63	337	16			20	030	804
US	20052	6717	71		<b>A1</b>	2	2005	1201	τ	JS	2005	-11	094	13			20	0504	121
PRIORIT	Y APPL	N. ]	INFO	. :					τ	JS	1992	-93	498	34		В2	19	920	325
									V	OV	1993	-US	781	4		A2	19	9308	324
									τ	JS	1994	-20	482	27		A2	19	9403	302
									F	ΞP	1993	-92	371	.4		А3	19	9308	324
									Ţ	JS	1993	-11	091	.1		A2	19	9308	324
									J	JS	1994	-29	446	8		<b>A</b> 1	19	9408	323
									τ	JS	1999	-28	808	0		A1	19	9904	108
									τ	JS	2001	-79	825	55		A1	20	0103	305
									ζ	JS	2002	-15	701	.9		A1	20	0205	530
									τ	JS	2003	-63	337	6		A1	20	0308	304

OTHER SOURCE(S): MARPAT 131:295568

AB  $\alpha$ - And  $\beta$ -Amino acid hydroxyethylamino sulfonamide compds. are effective as retroviral protease inhibitors, and in particular as inhibitors of HIV protease, as well as effective in preventing the growth of retroviruses in a solution General and specific schemes for chemical synthesis of the sulfonamide-containing hydroxyethylamine inhibitor compds. are described. Seventy-eight such compds. were tested for cytotoxicity and antiviral efficacy (IC50, EC50, and TD50 values at the nanomolar level are tabulated).

IT 169281-16-7P 247047-39-8P 247047-40-1P 247047-42-3P

RL: SPN (Synthetic preparation); PREP (Preparation) ( $\alpha$ - and  $\beta$ -amino acid hydroxyethylamino sulfonamides useful as retroviral protease inhibitors)

RN 169281-16-7 CAPLUS

CN Carbamic acid, [(1S,2R)-2-hydroxy-3-[[3-(methylthio)propyl](phenylsulfonyl)amino]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 247047-39-8 CAPLUS

CN Carbamic acid, [(1S,2R)-2-hydroxy-3-[(4-hydroxybutyl)(phenylsulfonyl)amino ]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

10/613,650

RN 247047-40-1 CAPLUS

CN Carbamic acid, [(1S,2R)-3-[[2-(dimethylamino)ethyl](phenylsulfonyl)amino]-2-hydroxy-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 247047-42-3 CAPLUS

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 12 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1998:799692 CAPLUS

DOCUMENT NUMBER:

130:38712

TITLE:

Preparation of  $\alpha$ - and  $\beta$ -amino acid

hydroxyethylamino sulfonamides useful as retroviral

protease inhibitors

10/613,650 Vazquez, Michael L.; Mueller, Richard A.; Talley, John INVENTOR (S): J.; Getman, Daniel; Decrescenzo, Gary A.; Freskos, John N. PATENT ASSIGNEE(S): G.D. Searle and Co., USA U.S., 67 pp., Cont.-in-part of U.S. Ser. No. 934,984, SOURCE: abandoned. CODEN: USXXAM DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE DATE APPLICATION NO. DATE PATENT NO. A 19981201 A2 19971203 A3 19981202 B1 20020605 US 1993-110911 19930824 EP 1997-113434 19930824 US 5843946 EP 810209 EP 810209 EP 810209

					2002000				
	R: AT,	BE,	CH,	DE,	DK, ES, FR,	GB, GR, IT, LI, LU,	NL,	SE, PT,	IE
AT	172717			E	19981115	AT 1993-923714		199308	24
ES	2123065			Т3	19990101	AT 1993-923714 ES 1993-923714		199308	24
AT	218541			E	20020615	AT 1997-113434 PT 1997-113434 ES 1997-113434		199308	24
PT	810209			${f T}$	20020930	PT 1997-113434		199308	24
ES	2177868			Т3	20021216	ES 1997-113434		199308	24
WO	9506030			A1	19950302	WO 1994-US9139		199408	
	W: AM,	AT,	AU,	BB,		CA, CH, CN, CZ, DE,			
	GE,	HU,	JP.	KE.	KG, KP, KR,	KZ, LK, LT, LU, LV,	MD.	MG MN	MW
	NL,	NO,	NZ,	PL.	PT, RO, RU.	SD, SE, SI, SK, TJ,	TT.	IIA IIS	UZ VN
	RW: KE,	MW,	SD,	AT.	BE, CH, DE,	DK, ES, FR, GB, GR,	TE.	TT LII	MC
	NL,	PT,	SE,	BF.	BJ. CF. CG.	CI, CM, GA, GN, ML,	MR.	NE SN	יים, דים דים
AU	9476697	•	•	A1	19950321	AII 1994-76697		100/00	າລໍ
	715618			A1	19960612	EP 1994-927162		199408	23
	715618			B1	19981216	21 1991 32,102		100400	23
		BE.	CH.		DK. ES. FR.	GB, GR, IE, IT, LI,	T.IT	NI. DT	CF.
АТ	174587	,	<b></b> ,	E	19990115	AT 1994-927162	ц,	199408	
	2127938			E T3	19990501	ES 1994-927162		199400	
	9500650			A	19950214	FI 1995-650		199502	
	112471			B1	20031215			199302	14
	5786483			A				100506	07
	5830897			A A	19981103			199506	07
	6172082			B1	20010109			199506	07
	5744481			A	19980428				
	6248775			B1	20010619			199704 199904	08
	6335460			B1	20020101	US 2000-510189		200003	22
	6472407			B1	20021029	US 2000-511005		200002	22
	6534493			B1	20030318			200012	21
	200205239	99		A1	20020502	US 2001-798255		200010	24 05
US	6417387			B2	20020709	00 2002 / 50055		200103	0.5
	200319131	L 9		A1	20031009	US 2002-157019		200205	3.0
US	6646010			B2	20031111	00 2002 137013		200203	50
	6924286			В1	20050802	US 2003-633376		200308	Λ4
PRIORITY	APPLN.	NFO.	:			US 2003-633376 US 1992-934984	ī	300300 32 199208	25
						EP 1993-923714		A3 199308	
						US 1993-110911		A 199308	
						WO 1993-US7814		A2 199308	
						US 1994-204827		A 199403	
						US 1994-294468		199408	
						WO 1994-US9139		N 199408	
							7	1 10050 <i>e</i>	43 07
						US 1995-485524	ī	A1 199506 B1 199506	07
						US 1999-288080	-	199506	0 /
								1 199904	
						US 2001-798255	,		
	•					US 2002-157019	F	1 200205	3 U

OTHER SOURCE(S):

MARPAT 130:38712

Amino acid hydroxyethylamino sulfonamide compds. P1NHCHR2CH(OH)CH2NR3SO2R4 [P1 = alkoxycarbonyl, aralkoxycarbonyl, alkanoyl, cycloalkylakoxycarbonyl, cycloalkylalkanoyl, aralkanoyl, aroyl, aryloxycarbonyl, heterocyclylcarbonyl, heterocyclyloxycarbonyl, heterocyclylalkoxycarbonyl, heteroaralkoxycarbonyl, heteroaryloxycarbonyl, heteroaroyl; R2 = alkyl, aryl, cycloalkyl, cycloalkylalkyl, (un)substituted aralkyl; R3 = H, alkyl, alkenyl, alkynyl, hydroxyalkyl, alkoxyalkyl, cycloalkyl, cycloalkylalkyl, heterocyclyl, heteroaryl, heterocyclylalkyl, aryl, aralkyl, heteroaralkyl; R4 = alkyl, haloalkyl, alkenyl, alkynyl, cycloalkyl, heterocycloalkyl, heteroaryl, aryl, aralkyl) were preparation as retroviral protease inhibitors. Thus,

N-[2R-hydroxy-3-[[(4-

methoxyphenyl)sulfonyl](2-methylpropyl)amino]-1S-(phenylmethyl)propyl]-4-pyridinecarboxamide was prepared by amidation of isonicotinoyl chloride hydrochloride with 2R-hydroxy-3-[(2-methylpropyl)](4-methoxyphenyl)sulfonyl]amino]-1S-(phenylmethyl)propylamine. Protease inhibitory data are tabulated.

IT 169281-15-6P 169281-16-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of amino acid hydroxyethylamino sulfonamides useful as retroviral protease inhibitors)

RN 169281-15-6 CAPLUS

CN Carbamic acid, [(1S,2R)-2-hydroxy-3-[(3-methoxypropyl)(phenylsulfonyl)amin o]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 169281-16-7 CAPLUS

CN Carbamic acid, [(1S,2R)-2-hydroxy-3-[[3-(methylthio)propyl](phenylsulfonyl)amino]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 13 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1998:157421 CAPLUS

DOCUMENT NUMBER:

128:204795

TITLE:

Preparation of THF-containing sulfonamides as

inhibitors of aspartyl protease

INVENTOR(S):

Tung, Roger D.

PATENT ASSIGNEE(S):

Vertex Pharmaceuticals Inc., USA

SOURCE:

U.S., 30 pp., Cont.-in-part of U.S. Ser. No. 393,460,

abandoned.

CODEN: USXXAM

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 5

	TENT NO.			KIN		APPLICATION NO.		
	5723490					US 1995-424819 EP 1998-113921		
EP	885887			A2	19981223	EP 1998-113921	19930907	
EP	885887			<b>A</b> 3	19990203			
EP	885887			B1	20030528			
	R: AT	, BE,	CH,	DE,	DK, ES, FR,	GB, GR, IT, LI, LU,	NL, SE, MC, PT, IE	;
	5585397			Α	19961217	US 1993-142327	19931124	
US	5783701			Α	19980721	US 1995-393460 CA 1996-2217737	19950223	
CA	2217737			AA	19961024	CA 1996-2217737	19960418	
WO	9633184					WO 1996-US5475	19960418	
	W: AL	, AM,	ΑT,	AU,	AZ, BB, BG,	BR, BY, CA, CH, CN,	CZ, DE, DK, EE,	
						KE, KG, KP, KR, KZ,		
	LU	, LV,	MD,	MG,	MK, MN, MW,	MX, NO, NZ, PL, PT,	RO, RU, SD, SE,	
		, SI			_			
	RW: KE	, LS,	MW,	SD,	SZ, UG, AT,	BE, CH, DE, DK, ES,	FI, FR, GB, GR,	
			LU,			BF, BJ, CF, CG, CI,		
	9655596			A1	19961107	AU 1996-55596	19960418	
	706732			B2	19990624	CN 1996-193364		
	1181755			A	19980513	CN 1996-193364	19960418	
	846110					EP 1996-912942	19960418	
EP	846110	D.E.	OI I		20020828			
	R: AT	, BE, , SI,	CH,	DE,	DK, ES, FR,	GB, GR, IT, LI, LU,	NL, SE, MC, PT,	
.TD	1050973		ы,	ъv, Т2		ID 1006 531054	10000410	
	3046357				19980922 20000529	JP 1996-531954	19960418	
	9608032			7	19990112	BR 1996-8032	10060410	
	306903			Δ	19990112 20000228	NZ 1996-306903	19960418 19960418	
	950			A	20010328	AP 1997-1119		
		. MW.	KE.		SD, SZ	AL 1997-1119	19900418	
AΤ	222761	,,	,	E	20020915	AT 1996-912942	19960418	
	291054			B6			19960418	
PT	846110			T	20021231		19960418	
	2181882			Т3	20030301	ES 1996-912942	19960418	
EE	4307			ום	20040616	PF 1007 266	19960418	
RO	119302			B1	20040013	RO 1997-1926	19960418 19960418	
SK	284785			В6	20051103	SK 1997-1431	19960418	
NO	9704722			Α	19971013	NO 1997-4722		
	317734			B1	20041213			
	63677			B1	20020930	BG 1997-102048	19971117	
RITY	APPLN.	INFO	.:			BG 1997-102048 US 1992-941982	B2 19920908	
						IIC 1992_1/2227	72 10021124	
						US 1995-393460 FB 1993-921428	B2 19950223	
						DF 1333-321440	A3 13330307	
						WO 1993-US8458	W 19930907	

US 1995-424819 A 19950419 WO 1996-US5475 W 19960418

OTHER SOURCE(S):

MARPAT 128:204795

GI

THF-containing sulfonamides (THF)R1NHCHDCH(OH)CH2ND'SO2E [I, R1 = CO, SO2, COCO, etc.; D, D' = aryl, carbocyclyl, heterocyclyl, alkyl, alkenyl; E = alkenyl, Het, O(Het), (Het) (Het), etc. with Het = carbocyclyl, aryl, heterocyclyl], which are aspartyl protease inhibitors, were prepared. E.g., epoxide II was treated with isobutylamine, 4-FC6H4SO2Cl, then deprotected and treated with N-succinimidyl-(S)-3-tetrahydrofuranyl carbonate to give a THF-containing sulfonamide. I are particularly well suited for inhibiting HIV-1 and HIV-2 protease activity and consequently, may be advantageously used as anti-viral agents against the HIV-1 and HIV-2 viruses.

IT 184357-39-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of THF-containing sulfonamides as inhibitors of aspartyl protease)

RN 184357-39-9 CAPLUS

CN Carbamic acid, [2-hydroxy-3-[(3-hydroxy-2,2-dimethylpropyl)][(4-methoxyphenyl)sulfonyl]amino]-1-(phenylmethyl)propyl]-, 1,1-dimethylethylester, [R-(R\*,S\*)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 79 THERE ARE 79 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1997:5844 CAPLUS

DOCUMENT NUMBER:

126:31265

TITLE:

Preparation of tetrahydrofuran-containing sulfonamide inhibitors of aspartyl protease for treatment of HIV

infection.

INVENTOR(S):

Tung, Roger D.

PATENT ASSIGNEE(S):

Vertex Pharmaceuticals Incorporated, USA

SOURCE:

PCT Int. Appl., 105 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE -------------------------WO 1996-US5475 WO 9633184 A1 19961024 19960418 W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML US 5723490 19980303 US 1995-424819 Α 19950419 AU 9655596 **A1** 19961107 AU 1996-55596 19960418 AU 706732 B2 19990624 EP 846110 **A1** 19980610 EP 1996-912942 19960418 EP 846110 B1 20020828 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI JP 10509739 T2 19980922 JP 1996-531954 19960418 JP 3046357 B2 20000529 BR 9608032 Α 19990112 BR 1996-8032 19960418 NZ 306903 Α 20000228 NZ 1996-306903 19960418 AT 222761 Ε AT 1996-912942 20020915 19960418 EE 4307 В1 20040615 EE 1997-266 19960418 RO 119302 В1 20040730 RO 1997-1926 19960418 SK 284785 В6 20051103 SK 1997-1431 19960418 NO 9704722 Α 19971013 NO 1997-4722 19971013 NO 317734 В1 20041213 BG 63677 B1 20020930 BG 1997-102048 19971117 PRIORITY APPLN. INFO.: US 1995-424819 A 19950419 US 1992-941982 B2 19920908 US 1993-142327 A2 19931124 US 1995-393460 B2 19950223 WO 1996-US5475 W 19960418

OTHER SOURCE(S): GT

MARPAT 126:31265

AB R1QNHCHR2CH(OH)CH2NR3SO2E [R1 = tetrahydrofuryl; Q = CO, SO2, COCO, O2C, OSO2, iminosulfonyl, aminocarbonyl, etc.; R2, R3 = (substituted) alkyl, alkenyl, carbocyclyl, cycloalkenyl, aryl, heterocyclyl; E = (substituted) heterocyclyl, carbocyclyl, aryl, heterocyclyloxy, carbocyclyloxy, aryloxy, amino, alkoxy, alkenyloxy, etc.], were prepared Thus, title compound (I), prepared from epoxide (II), showed Ki <0.1 nM against HIV-1 protease. IT 184357-39-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of tetrahydrofuran-containing sulfonamide inhibitors of aspartyl

protease for treatment of HIV infection)

RN 184357-39-9 CAPLUS

CN Carbamic acid, [2-hydroxy-3-[(3-hydroxy-2,2-dimethylpropyl)] (4-methoxyphenyl)sulfonyl]amino]-1-(phenylmethyl)propyl]-, 1,1-dimethylethylester, [R-(R\*,S\*)]- (9CI) (CA INDEX NAME)

## Absolute stereochemistry.

L4 ANSWER 15 OF 15 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:871984 CAPLUS

DOCUMENT NUMBER: 123:279761

TITLE: Hydroxyethylamino sulfonamides useful as retroviral

protease inhibitors

INVENTOR(S): Vazquez, Michael L.; Mueller, Richard A.; Talley, John

J.; Getman, Daniel P.; Decrescenzo, Gary A.; Freskos, John N.; Bertenshaw, Deborah E.; Heintz, Robert M.

PATENT ASSIGNEE(S): G.D. Searle and Co., USA; Monsanto Co.

SOURCE: PCT Int. Appl., 255 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 6

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9506030		WO 1994-US9139	19940823
		CA, CH, CN, CZ, DE,	
NL, NO, NZ	, RE, RG, RP, RR, , PL, PT, RO, RU,	KZ, LK, LT, LU, LV, SD, SE, SI, SK, TJ,	TT, UA, US, UZ, VN
RW: KE, MW, SD	, AT, BE, CH, DE,	DK, ES, FR, GB, GR,	IE, IT, LU, MC,
ND, PI, SE US 5843946	, Br, BU, Cr, CG, 10091201	CI, CM, GA, GN, ML, US 1993-110911	MR, NE, SN, TD, TG
119 6060476	7 20000500	US 1994-204827	19930824
33 0486608	A 20000509	05 1994-204827	19940302
AU 9476697	A1 19950321	AU 1994-76697	19940823
EP 715618	A1 19960612	EP 1994-927162	19940823
EP 715618	B1 19981216		
R: AT, BE, CH	, DE, DK, ES, FR,	GB, GR, IE, IT, LI,	LU, NL, PT, SE
US 6046190	A 20000404	US 1996-586866	19960124
PRIORITY APPLN. INFO.:		US 1993-110911	A 19930824
		US 1994-204827	A 19940302
		US 1992-934984	B2 19920825
		WO 1993-US7814	
		US 1994-204872	
		WO 1994-US9139	W 19940823
OTHER SOURCE(S):	MARPAT 123:2797	61	

AB Hyroxethylamino sulfonamide compds. AC(:Y)NR6CHR2CHOHCH2NR3S(:O)xR4 [I:

R2=(substituted)alkyl, aryl, cycloalkyl, cycloalkylalkyl, aralkyl; R3=H; R3,R4=R2, alkenyl, alkynyl, heterocycloalkyl, -aryl, -aralkyl, -cycloalkyalkyl; R6=H, alkyl; x=1,2; Y=O, S; A=RO, R; R=alkyl, alkenyl; (hetero)aryl, cycloalkyl, cycloalkylalkyl, aralkyl, NH2, mono- or disubstituted amino, etc.] are effective as retroviral protease inhibitors, and in particular as inhibitors of HIV protease. Many inhibitors were prepared by (1) preparing an N-protected amino epoxide and (2) reacting this with an amine and (3) preparing a sulfonamide by reacting with a sulfonyl chloride or sulfonyl anhydride in the presence of an acid scavenger. The amino function of the sulfonamide was then (4) deprotected and (5) reacted with a carboxylate. In vitro HIV protease assays with these compds. revealed inhibitors with IC50's as low as 1.4 nM, e.g. [1S-[1R\*(S\*),2S\*]]-I (A=p-MeOC6H4CH2OCONHCH2CHMe; Y=O; R6=H; R2=benzyl; R3=3-methylbutyl; x=2; R4=phenyl).

IT 169281-15-6P 169281-16-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(hydroxyethylamino sulfonamides useful as retroviral protease inhibitors)

RN 169281-15-6 CAPLUS

CN Carbamic acid, [(1S,2R)-2-hydroxy-3-[(3-methoxypropyl) (phenylsulfonyl)amin o]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 169281-16-7 CAPLUS

CN Carbamic acid, [(1S,2R)-2-hydroxy-3-[[3-(methylthio)propyl] (phenylsulfonyl )amino]-1-(phenylmethyl)propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

10/613,650

(FILE 'HOME' ENTERED AT 10:09:35 ON 23 MAY 2006)

FILE 'REGISTRY' ENTERED AT 10:09:41 ON 23 MAY 2006

L1 STRUCTURE UPLOADED

L2 4 S L1

L3 47 S L1 FULL

FILE 'CAPLUS' ENTERED AT 10:10:17 ON 23 MAY 2006 L4 15 S L3

=> d l1

L1 HAS NO ANSWERS

L1 STR

G1 C,S G2 O,S,N,P,CN,[@1],[@2]

Structure attributes must be viewed using STN Express query preparation.

=>

10/613,650

=> d his

(FILE 'HOME' ENTERED AT 10:12:56 ON 23 MAY 2006)

FILE 'REGISTRY' ENTERED AT 10:13:07 ON 23 MAY 2006

L1 STRUCTURE UPLOADED

L2 0 S L1

L3 0 S L1 FULL

=> d 11

L1 HAS NO ANSWERS

L1 STR

G1 C,S G2 O,S,N,P,CN,[@1],[@2]

Structure attributes must be viewed using STN Express query preparation.

=>